Photosensitized Electron Injection in Colloidal Semiconductors

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Abstract: Photosensitized electron injection in semiconductor particles was studied by using aqueous eosin (EO)/colloidal TiO₂ as the model system. Adsorption of EO onto the surface of TiO₂ occurs quantitatively at pH < 6 and leads to red-shifted absorption and emission spectra. Electron injection in the conduction band of the TiO₂ particles takes place from the S₀ state of adsorbed eosin and occurs with a rate constant of 8.5 × 10⁻¹⁰ s⁻¹ and a quantum yield of 38% (pH 3). The quantum yield decreases at high eosin occupancy of the particles due to concentration quenching. Back-electron transfer occurs via a rapid intraparticle reaction between EO⁺...e⁻ pairs associated with the same TiO₂ host aggregate and via a slower process involving bulk diffusion. The rate constant for intraparticle recombination is 2 × 10⁻⁵ s⁻¹, i.e., 4 × 10⁶ times slower than that for electron injection. This enables light-induced charge separation to be sustained on a colloidal TiO₂ particle for several microseconds, which is sufficient to trap the electron by a noble metal deposit. Implications for light-energy conversion devices are discussed.

The photosensitization of electron transfer across the semiconductor solution interface plays a vital role in silver halide photography1 and electrophotography.2 Recently, it has also gained interest with regards to light energy conversion in photoelectrochemical cells.3 Effort in this area has concentrated on improving the visible light response of wide-band semiconductor materials such as ZnO and TiO₂. Sensitization is achieved by adsorption of dye molecules at the semiconductor surface which, upon excitation, inject an electron into its conduction band. The first successful experiment of this type was described by Putzeiko and Terenin,4 who found that the Dember effect of ZnO powder in visible light was sensitized by xanthene and cyanine dyes. Numerous investigations with ZnO have followed their original report, owing mainly to the successful application of this material in reprographic processes.5 Sensitization processes investigated with TiO₂ comprise the reduction of O₂ by hydroquinone6 and water cleavage7 as well as hydrogen generation8 by visible light. Phthalocyanines,9 Ru(bpy)₃⁺, and derivatives10 as well as a Ti₄⁺-chelate with 8-hydroxyquinolines11 were employed as sensitizers.

While much important knowledge has been gathered over the years on the overall performance of dye-sensitized semiconductor systems,12 more information about the details of the electron-injection process is urgently required. The rapid nature of these reactions requires application of fast kinetic techniques which in the case of solid electrodes or powders is very difficult. On the other hand, semiconductor particles of colloidal dimensions are sufficiently small to yield transparent solutions, allowing for direct analysis of interfacial charge-transfer processes by a laser photolysis technique.10 So far, there are very few time-resolved studies of dye sensitization with colloidal semiconductor particles. Terenin and Akimov11 allude to flash photolysis of AgI colloids in the presence of dyes. Kiwi12 examined electron transfer from excited Ru(bpy)₃⁺ to colloidal TiO₂ at elevated temperature. Very recently, the erythrosine sensitization of colloidal TiO₂ in acetonitrile has been studied by Kamat and Fox.13

The present study employs eosin Y as a sensitizer to study electron injection in colloidal TiO₂ particles. Precise information is available on the ground- and excited-state properties of this dye as well as the absorption spectra of its monoeclorinated and monoxygenated radical forms, facilitating identification of transient species formed during the photoreaction. Apart from the initial charge-transfer event, we explore the fate of the injected electron, in particular its back reaction with the cation radical of the sensitizer, and the competing trapping by noble metals deposited on the surface of the semiconductor particle.

Experimental Section

Materials. TiCl₄ (Fluka purissimum) was further purified by vacuum distillation (40 °C, ca. 25 torr) to yield a colorless liquid. Colloidal TiO₂ particles were prepared by hydrolysis of this material at 0 °C according to a procedure previously described.13 The mean particle radius obtained from electron microscopy and quasi-elastic light scattering is 50–60 Å. Electrophotoretic measurements give a point of zero z potential of 4.7. At pH ≤ 4 and pH ≥ 9, the TiO₂ sol is stable over at least several days, no

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The absorption spectra were recorded on a Cary 219 (Varian) spectrophotometer. Significant aggregation being detectable by a quasi-elastic light-scattering technique. In the intermediate pH domain the colloid requires a protective agent to prevent flocculation. Poly(vinyl) alcohol was used in this case to stabilize the colloidal particles. Commercial PVA (Mowiol 98–10, Hoechst, West Germany) was treated by UV light to remove impurities. Control experiments showed that the protective agent had practically no effect on the characteristics of sensitized electron injection.

Eosin Y, disodium salt (Fluka) was purified by column chromatography with use of neutral A13O3. After it is eluted with a mixture of methanol/water/triethanolamine (90/9/1, v/v/v), the principal fraction is acidified with HCl and the precipitate is filtered and washed. The protonated form of eosin is finally recrystallized from water until the product showed in alkaline solution an extinction coefficient for the protonated form of the monomer dye. Decreasing the pH below 5 shifts the absorption band to the red with \( \lambda_{\text{max}} \) 520 nm at pH 3. We attribute this to the protonation of the phenolic hydroxyl group. Acid–base titration revealed the presence of one equivalent point, indicating \( pK_a \approx 4.4 \). The second \( pK_a \) could not be determined by this method since the neutral eosin molecule is insoluble in water.

Figure 1 shows the effect of \( pK_a \) on the eosin absorption in the presence of TiO2 (0.5 g/L). At pH 10 the spectrum is practically identical with that found in neutral or basic water. Decreasing the pH from 7.5 to 5 shifts the maximum to 527 nm. Associated with this transition is an isosbestic point at 523 nm. Within the pH range from 5 to 3 the spectral features remain essentially constant. The observed red shift in the absorption spectrum without notable change in its shape can be unambiguously attributed to the adsorption of monomer eosin dianion on the colloidal TiO2 particles. Coulombic interaction with the TiO2 surface appears to play an important role since the spectral transition in Figure 1 occurs in a pH domain where the charge of the colloid changes from negative to positive. Quantitative determination of the adsorbed amount of eosin was carried out with 20 mL of buffered solution containing colloidal TiO2 (2.5 g/L), 5 x 10⁻⁵ M eosin, and PVA stabilizer (5 g/L). The solution is enclosed in a dialysis membrane (cellulose acetate, pore diameter 15–30 Å) and stirred in 80 mL of buffered aqueous ambiant. The equilibrium was reached after ca. 40 h, and the concentration of free eosin was determined spectrophotometrically. From this procedure we find for the adsorbed fraction of eosin at pH 10, 7.2, and 4 the values 0.08, 0.52, and 0.95, respectively. This confirms the assignment of the spectra obtained for \( pK_a \leq 5 \) in the presence of TiO2 (0.5 g/L) protected by PVA (1 g/L).

Results and Discussion

(i) Effect of Colloidal TiO2 on the Absorption and Emission Characteristics of Eosin. The absorption spectrum of a 10⁻⁵ M eosin solution in neutral or basic water shows a characteristic maximum at 516 nm with \( \varepsilon = 9.7 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1} \) which arises from the dianionic form of the monomer dye. Decreasing the pH below 5 shifts the absorption band to the red with \( \lambda_{\text{max}} \) 520 nm at pH 3. We attribute this to the protonation of the phenolic hydroxyl group. Acid–base titration revealed the presence of one equivalent point, indicating \( pK_a \approx 4.4 \). The second \( pK_a \) could not be determined by this method since the neutral eosin molecule is insoluble in water.

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(19) Eosin Y has only a weak tendency to form dimeric aggregates in aqueous solution (dimerization equilibrium constant \( 9 \times 10^{-10} \)). Spectral evidence for dimer formation (\( \lambda_{\text{max}} \) 531 and 487 nm, \( c_{\text{eq}} \) 40500 and 51000 M⁻¹ cm⁻¹) was found only at dye concentrations exceeding \( 5 \times 10^{-5} \) M.


(21) From spectral analysis we estimate for the protonation of the carboxyl group \( pK_a \approx 3.4 \).

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Figure 3. Effect of pH on the luminescence quantum yield of eosin in the presence of TiO₂ (500 mg/L); φ₀ values obtained from integration of the emission curves in Figure 2. The quantum yield in pure water φ₀ = 0.22 was taken from Soep et al.27

Figure 4. Effect of TiO₂ concentration on the eosin emission spectrum at pH 4; dye concentration 10⁻⁵ M. Practically identical results were obtained for unprotected and PVA-protected TiO₂ solutions.

Figure 1 to the eosin associated with the surface of TiO₂ particles.

Figure 2 illustrates the effect of pH on the luminescence of eosin in the presence of colloidal TiO₂ (500 mg/L). At pH 10 the emission maximum is located at 540 nm and is identical with that found in water. Decreasing the pH from 7 to 5.2 leads to an abrupt red shift in the emission and marked reduction in its intensity. Below pH 5.2 the maximum in the luminescence remains at 550 nm while the intensity is further decreased. Figure 3 compares fluorescence quantum yields at the excitation wavelength (A 500 nm) for aqueous eosin solution of different pH in the presence of TiO₂ (500 mg/L). The quantum yield for the TiO₂ solution drops sharply between pH 6 and 4, i.e., in a pH domain where the transition from the free to the surface-bound state of the eosin dianion occurs. Since in TiO₂-free solution the fluorescence changes very little between pH 7 and 4, one infers from these results that the luminescence of eosin is strongly quenched by the TiO₂ particles.

In Figures 4 and 5 the pH of the aqueous solution was maintained at 4 and the effect of colloidal TiO₂ on the eosin luminescence was investigated. Addition of TiO₂ (50 mg/L) leads to a pronounced decrease in the luminescence yield (φ₀) and to a red shift in the emission maximum (λ_max). With 100 mg/L of TiO₂ present, φ₀ has dropped to 14% of its initial value and λ_max is ca. 548 nm. At this TiO₂ concentration φ₀ passes through a minimum. Further increase in TiO₂ concentration enhances the luminescence which at [TiO₂] ≈ 0.5 g/L reaches a plateau corresponding to 32% of its intensity in pure water. No changes in λ_max occur in this concentration domain.

Since at a TiO₂ concentration of 100 mg/L eosin is quantitatively associated with the colloidal particles, the emission with λ_max 548 nm can be attributed to the adsorbed state of the dye. Red shifts in the emission of xanthene dyes upon adsorption to metal oxide surfaces have been previously observed19 and are due to hydrogen bond formation with surface hydroxyl groups. The displacement of the maximum is related to the polarizability and electronegativity of the metal cation. Thus, for fluorescein adsorbed onto Zn(OH)₂ and Al(OH)₃, Δλ_max is 150 and 30 nm, respectively.25 The relatively small red shift observed here for eosin indicates binding to OH groups at the TiO₂ surface with relatively acidic character.

In order to rationalize the drastic reduction in the eosin luminescence intensity upon adsorption to the TiO₂ particles observed in Figures 4 and 5, we consider first concentration quenching. Deactivation via such a mechanism is suggested by the observation in Table I that the smallest emission intensity is obtained for the highest mean occupancy of the TiO₂ particles by eosin, i.e., 160 molecules/particle. Under these conditions the surface concentration of eosin is 2.1 × 10⁻⁵ cm⁻² and the mean distance between adjacent dye molecules is 22 Å. We expect in this case efficient dipolar energy transfer between adjacent eosin molecules until trapping by a molecular configuration favorable for radiationless deactivation quenches the excited state.24 Concentration quenching was also encountered by Liang et al. in very recent

Table I. Concentration of TiO₂ Particles and Average Occupancy of the Particles by Eosin at 10⁻⁵ M Dye Concentration as a Function of Total TiO₂ Concentration

<table>
<thead>
<tr>
<th>[TiO₂], g/L</th>
<th>[particles], mol L⁻¹</th>
<th>av. no. of eosin molecules per particle</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.1</td>
<td>6.3 × 10⁻⁸</td>
<td>160</td>
</tr>
<tr>
<td>0.2</td>
<td>1.3 × 10⁻⁷</td>
<td>80</td>
</tr>
<tr>
<td>0.5</td>
<td>3.1 × 10⁻⁷</td>
<td>32</td>
</tr>
<tr>
<td>1.0</td>
<td>6.3 × 10⁻⁷</td>
<td>16</td>
</tr>
<tr>
<td>3.0</td>
<td>1.9 × 10⁻⁵</td>
<td>5</td>
</tr>
</tbody>
</table>

(24) We shall publish a detailed treatment of concentration quenching involving eosin absorbed onto colloidal TiO₂ particles elsewhere. (Rothenberger, G.; Humphry-Baker, R.; Moser, J.; Grätzel, M., manuscript in preparation.)
monitored at 600 nm where there is little interference from the triplet reaction. From the kinetic analysis, we obtain for the quantum yield of 0.76,26,27 Figure 6. In deaerated solution the ground-state absorption recorded at 530 nm.

The end of the pulse spectrum is identical with that of the triplet state. These authors concluded that the reduction was due only to concentration quenching and not electron injection from the excited dye in the semiconductor. Consistent with the deactivation of excited eosin via concentration quenching is the observation in Figure 5 that the emission quantum yield passes through a minimum at 100 mg of TiO$_2$/L, i.e., under conditions where the occupancy of TiO$_2$ particles by eosin is highest. Increasing the TiO$_2$ concentration decreases the particle occupancy by eosin (Table I), and this should diminish the efficiency of concentration quenching and not electron injection from the ground-state (EO(S$_0$)) which in aqueous solution is formed with a lifetime 95 ps.

(i) Laser Photolysis Studies of Sensitized Electron Injection: Effect of pH and TiO$_2$ Concentration. Laser photolysis of eosin was first performed in pure water (pH 4) in the absence of TiO$_2$. Transients were identified on the basis of earlier flash photolysis studies and literature absorption data for pertinent intermediates. The end of the pulse spectrum is identical with that of the triplet state (EO(T$_1$)) which in aqueous solution is formed with a quantum yield of 0.76,26,27 Figure 6. In deaerated solution the triplet decay obeys mixed first-/second-order kinetics. It is monitored at 600 nm where there is little interference from the absorption of ground-state eosin or ionic products formed during the triplet reaction. From the kinetic analysis, we obtain for the first half-lifetime 95 ps. Concomitantly with the triplet decay occurs the formation of eosin cation radicals (EO$^+$) monitored at 450 nm. The EO$^+$ concentration attains a maximum after ca. 200 μs and decreases subsequently according to a second-order rate law. Associated with this decay is the recovery of EO(S$_0$) ground-state absorption recorded at 530 nm.

In accordance with earlier propositions,27-29 we explain this behavior with the following scheme (literature rate constants are given below the respective equation)

$$EO(S_0) \xrightarrow{h} EO(S_1) \xrightarrow{\phi_{TS}} EO(T_0) \quad (2)$$

$$EO(T_1) \xrightarrow{\cdot} EO(S_0) \quad (3)$$

$$540 \text{ s}^{-1}, 700 \text{ s}^{-1} \quad (4a)$$

$$EO(T_1) + EO(S_0) \rightarrow 2EO(S_0) \quad (4b)$$

$$3 \times 10^4 \text{ M}^{-1} \text{ s}^{-1} \quad (4c)$$

$$EO(T_1) + EO(S_0) \rightarrow EO^+ + EO^- \quad (5a)$$

$$2EO(T_1) \rightarrow 2EO(S_0) \quad (5b)$$

$$2EO(T_1) \rightarrow EO^+ + EO^- \quad (5c)$$

$$\text{eq 5a and 5b: } 1.3 \times 10^9 \text{ M}^{-1} \text{ s}^{-1} \quad (6)$$

$$8.8 \times 10^7 \text{ M}^{-1} \text{ s}^{-1} \quad (6)$$

The rate constants determined in our study are in agreement with the literature results apart from the value of k$_4$ for which we find $5 \times 10^8 \text{ M}^{-1} \text{ s}^{-1}$. We confirm also that the formation of eosin cation radicals arises from the disproportionation of triplet states. (The concomitant formation of EO$^-$ could not be monitored since at pH 4 EO$^-$ is protonated, pK 6.5,23.) From the one-electron oxidation and reduction potentials of ground-state eosin, $E^0$ (EO/EO$^+$) = 1.1 V$^{28}$ and $E^0$ (EO/EO$^-$) = 0.8 V (NHE)$^{31}$ and the triplet-state energy of 1.98 eV$^{30}$, the driving force for reaction 6 is only ca. 0.1 eV. The contribution of this process to EO$^+$ formation is therefore small compared to that of reaction 5b.

Addition of colloidal TiO$_2$ introduces striking changes in the photoredox behavior of eosin. The transient spectrum obtained immediately after laser excitation of a deaerated $10^{-3}$ M eosin solution in water (pH 4) containing colloidal TiO$_2$ (500 mg/L) is shown in Figure 7. It is distinguished by a prominent peak at 475 nm, a much weaker and broad absorption above 500 nm, and a shoulder at 400 nm. The spectrum is corrected for ground-state absorption within the wavelength region 440 nm ≤ λ ≤ 570 nm. In Figure 7 the intense absorption with λ$_{max}$ 475 nm can be unambiguously attributed to the semioxidized eosin, EO$^+$. Spectral characteristics are identical with those observed.


(31) Literature values for $E^0$(EO/EO$^+$) are $-0.85^{26}$ and $-0.79$ V.$^{32}$

for EO⁺ in water except that \( \lambda_{\text{max}} \) is red shifted by 13 nm. A similar red shift, i.e., from 516 to 527 nm, occurred in the absorption maximum for EO(S0) in Figure 1 upon adsorption onto the TiO₂ particles and was attributed to hydrogen bond formation with surface hydroxyl groups. That EO⁺ is produced from photoexcitation of EO(S0) in colloidal TiO₂ solution was recently confirmed by Rossetti and Brus using time-resolved laser Raman spectroscopy. The Raman spectrum of EO⁺ associated with TiO₂ particles was different from that in pure water, and this was attributed to protonation of EO⁺ by surface hydroxyl groups. The effect of TiO₂ concentration on the quantum yield of EO⁺ formation, \( \phi(\text{EO}^+) = [\text{EO}^+]/[\text{EO}(S_0)] \), where \([\text{EO}(S_0)]\) is the concentration of eosin excited by the laser pulse, is illustrated in Figure 8. Concentrations of EO⁺ were calculated from the increase in 450-nm absorbance during the laser pulse by using \( \varepsilon_{450} \text{(EO⁺)} = 2.5 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1} \). The signal at 530 nm contains a negative contribution from EO(S₂), that in pure water, for the triplet state at 530 and 600 nm, the error limit of the \( \phi(\text{EO}^+) \) values is 10%.

In Figure 7 the broad absorption at 600 nm and the shoulder at 400 nm arise from eosin triplet state and eosin cation radical, respectively. The quantum yield for triplet formation is also affected by the TiO₂ concentration, and these results are included in Figure 8. These values were determined from the increase in 600-nm absorbance during the laser flash by using \( \varepsilon_{600} \text{(EO(T₁))} = 1.2 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1} \). Quantum yield for EO⁺ formation increases from 27% to 35% upon increasing the TiO₂ concentration from 0.1 to 0.5 g/L, and reaches a plateau value of 38% at 3 g/L of TiO₂.

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The effect of pH on the quantum yield of eosin cation radical and triplet formation at a fixed concentration of colloidal TiO₂ of 500 mg/L, [PVA] = 1 g/L.

It was shown above that in pure water the photogeneration of EO⁺ is a relatively slow and inefficient process, arising from the dissociation of eosin triplet states. By contrast, in the presence of colloidal TiO₂, EO⁺ is generated efficiently and at a high rate, its formation being completed within the 15-ns duration of the laser pulse. In this case the mechanism of EO⁺ formation is different from that in water and involves electron injection from the electronically excited eosin into the conduction band of the colloidal TiO₂ particles. Furthermore, we can identify the S₁ state of the dye as the active intermediate in the charge-transfer process, i.e.

\[
\text{EO}(S₁) \rightarrow \text{EO}^+ + \text{e}_{\text{CB}} (\text{TiO}_2) \tag{7}
\]

The thermodynamic driving force for reaction 7 is given by the difference between the reduct potential of EO(S₁) and the conduction band position of colloidal TiO₂, \( E_{\text{CB}} \). For the former a value of -1.2 V (NHE) is derived from the ground-state potential and the singlet excitation energy of 2.3 eV. There has been experimentally determined as \( \Delta G^\circ = -0.1 \text{ to } -0.059 \text{ pH} \). Thus, there is sufficient driving force for reaction 7 even under alkaline conditions. Nevertheless, charge injection is only observed at pH < 6, i.e., under conditions where eosin is associated with the TiO₂ particles. This is reasonable since close proxility of the reactants is required for electron transfer to compete with the two other channels of singlet deactivation, i.e., intersystem crossing as well as radiative and nonradiative deactivation.

\[
\text{EO}(S₁) \rightarrow \text{EO}^+ + \text{e}_{\text{CB}} (\text{TiO}_2) \tag{7}
\]

The formation of EO⁺ during the photolysis of eosin/TiO₂ solutions could arise from reactions other than (7). One possibility is electron transfer from the triplet state into the conduction band of the TiO₂ particles:

\[
\text{EO}(T₁) \rightarrow \text{EO}^+ + \text{e}_{\text{CB}} \tag{10}
\]

The driving force for reaction 10, derived from the standard potential of the EO(T₁)/EO⁺ couple, \( E^\circ = -0.9 \text{ V (NHE)} \), is ca. 0.5 eV at pH 5, increasing further by 59 mV with each decreasing pH unit. In spite of these favorable thermodynamic conditions, charge injection from the triplet does not occur. The triplet lifetime for eosin associated with colloidal TiO₂ is the same as that in pure water, i.e., between 100 and 200 ns. This shows that any electron transfer via reaction 10 must be much slower than the other triplet deactivation processes.

A second possibility is the generation of EO⁺ via triplet-ground state or triplet-triplet dismutation, reactions 4b and 5b. While

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**Figure 8.** Effect of TiO₂ concentration on the quantum yield of eosin cation radical and triplet formation. Conditions as in Figure 7.

**Figure 9.** Effect of pH on the quantum yield of eosin cation radical and triplet formation at a fixed concentration of colloidal TiO₂ of 500 mg/L, [PVA] = 1 g/L.
these processes are comparatively slow in pure water, they could be accelerated significantly when they occur on a TiO₂ particle. The kinetics of chemical reactions between species restricted in their diffusional motion to the surface of a spherical aggregate have been previously analyzed. For a pair of eosin triplets associated with a 110 Å size TiO₂ particle, the average time between two encounters is 0.07 and 1.5 μs depending on the diffusion coefficient of the reactants at the TiO₂ surface. This is much longer than the EO⁺ formation time which was observed here to occur within the 10-ns laser pulse. Note also that (EO⁺) in Figure 9 increases with TiO₂ concentration, i.e., with decreasing eosin occupancy of the particles. The opposite trend would be expected if intraparticle triplet dismutation made a significant contribution to EO⁺ formation. Furthermore, (EO⁺) remains at the maximum value of 38% at pH 4 even when only one eosin molecule is associated with a TiO₂ particle, i.e., under conditions where no intraparticle reactions are possible.

We thus conclude that the primary photochemical events in the eosin/TiO₂ solutions are adequately described by eq 7–9. The relative contributions of these three reactions to the deactivation of EO(S₁) depend strongly on both TiO₂ concentration and pH. In order to explain the TiO₂ concentration dependency of the parameters $\phi_f$, $\phi_{EO⁺}$, and $\phi_T$, we draw attention to the fact that the nonradiative decay constant, and hence $\tau = (k_{ir} + k_i + k_{irc} + k_{inj})^{-1}$

\[(11)\]

are strongly influenced by the eosin occupancy of the particles. The closer the distance between adjacent dye molecules on the TiO₂ surface, the more efficient will be the concentration quenching, i.e., the higher the value of $k_{ir}$. As a consequence, the parameters $\tau$, $\phi_f$, and $\phi_T$ increase and attain a plateau at higher TiO₂ concentration where the influence of concentration quenching becomes negligible. The effect of pH is to change the rate of charge injection apart from influencing the adsorption of eosin to the TiO₂ particles. Thus, in the pH domain from 5 to 2, practically all the eosin is associated with the TiO₂ colloid. Nevertheless the quantum yield of EO⁺ formation increased from 27% to 45%. This is attributed to an increase in the charge injection rate constant $k_{inj} = \phi_{EO⁺}/\tau$

\[(14)\]

resulting from the augmentation of the driving force for reaction 7 which is 0.8 and 0.98 eV at pH 5 and 2, respectively. Apparently, in the photosensitized electron transfer from eosin to the conduction band of TiO₂, a relatively large expenditure of free energy is required to obtain fast reaction rates. We have calculated the absolute value of $k_{ir}$ for 3 g/L TiO₂ solutions of pH 3 where $\phi_f = 0.09$, $\phi_T = 0.18$, and $\phi_{EO⁺} = 0.4$. Using eq. 11–14 we derive $k_i/k_{ir} = 0.27$, $k_{irc}/k_{ir} = 0.55$, and $k_{inj}/k_{ir} = 1.21$. Assuming that the radiative decay for eosin(S₁) adsorbed on TiO₂ occurs with the same rate as in ethanol, $k_i = 1.91 \times 10^{8}$ s⁻¹ (the emission spectrum of EO(S₁) in ethanol has the same shape and $\lambda_{max}$ as in colloidal TiO₂ solution), one obtains $k_{inj}(\text{pH } 3) = 8.5 \times 10^8$ s⁻¹


\[(39)\] Detailed analysis is given in ref 38 for the case of reactions on the surface of micellar aggregates. Since eosin is linked to the TiO₂ particles through hydrogen bonds, its diffusional motion on the surface may be slower than that considered there.

\[(40)\] We attribute this irreversible bleaching to a side reaction of EO⁺ which by analogy with fluorescein is probably dimerization, c.f.: Lindquist, L. J. Phys. Chem. 1963, 67, 1701.
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Figure 11. Schematic illustration of charge injection and intraparticle back-electron transfer in the photosensitization of a colloidal semiconductor particle: (a) without redox catalyst, (b) noble metal catalyst-loaded particle.

oxidized eosin behind on the surface. The subsequent back-electron transfer

\[ e_{CB}^- + EO^+ \rightarrow EO(S_0) \]  

occurs in two steps.

The fast initial part involves \( e_{CB}^- \ldots EO^+ \) pairs associated with their original TiO\(_2\) host particle. Such an intraparticle process is expected to obey first-order kinetics\(^{(41-43)}\) if only one pair per particle participates in the reaction, as is the case under the experimental conditions employed in Figure 10. Only a fraction of 50\% of the eosin cation radicals produced by the laser pulse undergoes this rapid back reaction. The remainder desorbs from the TiO\(_2\) particle and escapes into the solution bulk. This part will undergo slow back reaction which involves diffusional encounter of \( EO^+ \) and TiO\(_2\) particles (interparticle mechanism). For the rapidly decaying fraction of \( EO^+ \), we can write

\[ f(EO^+) = \frac{k_b}{k_b + k_d} = \frac{k_b}{k_{obsd}} = 0.5 \]  

where \( k_b \) and \( k_d \) are the specific rates (in s\(^{-1}\)) of intraparticle back reaction and desorption of \( EO^+ \) and \( k_{obsd} = 4 \times 10^3 \) s\(^{-1}\) is the observed rate constant for the fast \( EO^+ \) decay. From eq 16 we obtain

\[ k_b = 2 \times 10^5 \text{ s}^{-1} \]

corresponding to a mean lifetime of 5 \( \mu \text{s} \) for a \( e_{CB}^- \ldots EO^+ \) pair associated with a TiO\(_2\) particle. The intraparticle back reaction itself is composed of two elementary steps comprising diffusion of the injected charge to the particle surface and subsequent interfacial electron transfer to the \( EO^+ \) parent ion. The rate parameter \( k_b \) can therefore be expressed as

\[ k_b^{-1} = k_{diff}^{-1} + (k_{inj})^{-1} \]

(17)

where \( k_{diff} \) and \( k_{inj} \) are the specific rates of these individual processes. Since

\[ k_{diff} \approx \frac{1}{\tau_{diff}} \]

(18)

where \( \tau_{diff} = 2.5 \) ps is the average diffusion time for an electron from the particle interior to the surface and \( r_s = 3.6 \times 10^{-3} \) is the area ratio of an eosin molecule and the surface of a TiO\(_2\) particle, we can estimate for \( k_{diff} \) a value of \( 1.5 \times 10^9 \) s\(^{-1}\). This is much bigger than the experimentally derived value for \( k_b \). Therefore, in the back reaction of \( e_{CB}^- \) with \( EO^+ \), the electron transfer at the interface is rate determining, \( k_b \approx k_{inj} \). Note that this reverse electron transfer occurs at ca. \( 4 \times 10^3 \) times smaller rate than the forward injection from the EO(S\(_0\)) state into the TiO\(_2\) particle. Such a behavior is in accordance with the predictions of classical electron-transfer theory.\(^{(45)}\) The highly exothermic nature of the back-electron transfer (\( \Delta G = -1.5 \) eV) is expected to result in a decrease in reaction rate. However, we draw attention to the fact that classical theory may not be applicable in the inverted region. Thus, reduction of aromatic cation radicals by conduction band electrons of ZnO electrodes is not significantly affected by the driving force and occurs very rapidly despite the highly exothermic character of these reactions.\(^{(46)}\) (One referee pointed out that the entropy, i.e., phase space factor associated with the volume of the TiO\(_2\) particle, should make the back-transfer rate slow. Marcus theory does not incorporate this.)

The relatively slow rate of intraparticle back reaction between \( e_{CB}^- \) and \( EO^+ \) is a desirable feature from the viewpoint of light energy conversion. It allows photoinduced charge separation to be sustained over a sufficiently long time so that a significant fraction of \( EO^+ \) can escape from the TiO\(_2\) particles into the solution. Once in the aqueous bulk, \( EO^+ \) will still undergo back reaction with \( e_{CB}^- (\text{TiO}_{2}) \). However, this involves bulk diffusion and hence occurs at a much slower rate than intraparticle back-electron transfer. Desorption of \( EO^+ \) from the surface of TiO\(_2\) is more easy than that of the eosin parent molecule due to decreased Coulombic attraction.

(iv) Photosensitized Electron Injection into Noble Metal Loaded TiO\(_2\) Particles. In the following we report on some preliminary

\( (41) \) The situation is equivalent to the case of intramicellar electron-transfer kinetics which are dealt with in references 38 and 43.


experiments designed to study the trapping of photoinjected electrons by catalysts deposited on the surface of the TiO₂ particles. Figure 12b shows oscilloscope traces obtained from the laser photolysis of eosin in colloidal TiO₂ loaded with 0.5% RuO₂. The vertical rise of the absorbance signal at 470 nm indicates formation of EO⁺ during the light pulse. From this absorbance increase the quantum yield of charge injection was determined at 0.4 which is slightly higher than in the absence of RuO₂. The features of the EO⁺ decay are strikingly different from that observed for bare TiO₂ particles in that there is no fast component characteristic for intraparticle electron transfer. The EO⁺ absorption decays here in a smooth and relatively slow fashion with a first half-life of 240 as. Kinetics of ground-state bleaching monitored at 530 nm correspond to the mirror image of the 470-nm absorbance decay.

Similar effects are observed when Pt instead of RuO₂ is deposited onto the TiO₂ particles, Figure 12a. In this case, more catalyst is required to eliminate the fast step in the EO⁺ reaction. Thus, at 1% Pt loading the initial rapid decline in the 470-nm signal is still perceptible and constitutes 25% of the total absorbance change induced by the laser pulse, as compared to 50% for Pt-free TiO₂ particles.

These observations are rationalized in terms of the scheme shown in Figure 11b. The role of the catalyst is to trap photo-injected electrons and to intercept in this way their back reaction with EO⁺ adsorbed at the particle surface. This allows a larger fraction, or all of the EO⁺ formed during the laser pulse, to escape in solution. As discussed above, the subsequent back reaction is relatively slow since it involves bulk diffusion. Also, the species that reduced EO⁺ is not e⁻ but rather an H⁺ atom adsorbed at the catalyst surface, or in the case of RuO₂, a ruthenium lattice ion with a <4+ valency.

From Figure 11a one can derive an absolute value for the rate constant of electron transfer from the conduction band of the TiO₂ particle to the Pt deposit on the surface. At 1% loading the fraction of EO⁺ participating in the fast decay is 0.25, and the observed rate constant for this rapid charge recombination process is 6 × 10⁷ s⁻¹. By analogy with eq 16, we write

\[
\frac{f(EO^+)}{k_\text{obsd}} = 0.25
\]  

where

\[
k_\text{obsd} = k_b + k_d + k_\text{trasp}
\]

One obtains therefore \( k_\text{trasp} = 2.5 \times 10^6 \) s⁻¹.

A conspicuous feature of Figure 12 is that at comparable loading RuO₂ is more effective as a e⁻ scavenger than Pt. Trapping efficiencies are related to the dispersion of the catalyst and the nature of the junction formed at the semiconductor/metal interface. Ohmic contact is necessary for electron transfer to occur rapidly. However, from the work function of Pt (\( \phi = 5.2 \text{ eV} \)) or RuO₂ (\( \phi = 4.8 \text{ eV} \)) and the electron affinity of TiO₂ (\( x = 4.0 \text{ eV} \)), one would predict that a rectifying barrier of 1.2 or 0.8 V, respectively, should be produced at the junction of TiO₂ with these catalysts. This would impair electron transfer from the conduction band of the particle to the noble metal deposit. However, Hope and Bard⁵⁰ have shown that the electrical contact between Pt and TiO₂ is often ohmic due to the interdiffusion of the two materials. They also predict that due to the high density of surface states the junction between a colloidal TiO₂ particle and Pt should have a low resistance. This view is supported by the work of Aspnes and Heller,⁵¹ who found that hydrogen adsorption on Pt decreases its work function, rendering its contact with TiO₂ ohmic. The eosin/TiO₂ system lends itself to an investigation of the effects of ambient such as H₂ on the rate of electron trapping by a noble metal deposit.⁵² Such studies are expected to give clues as to the nature of the catalyst/TiO₂ junction when the particles are suspended in water and exposed to reducing or oxidizing atmospheres. These effects are presently being investigated further.

Conclusions

Using eosin/colloidal TiO₂ solutions as a model system, we have performed here a detailed study to scrutinize the salient features that control photosensitized electron injection from an excited dye in the conduction band of a semiconductor particle. Charge transfer occurs from the excited eosin singlet and competes with other radiative and nonradiative deactivation routes. Adsorption of eosin to the TiO₂ surface is a prerequisite for successful photosensitization. Low pH and high TiO₂ concentration favor charge injection. This is due to an increase in driving force of the reaction and dilution of the surface concentration of the dye, respectively. At high eosin occupancy of the particles, concentration quenching occurs, and we shall model this process using a statistical approach that takes into account the distribution of interacting species in the compartmentalized reaction space. Concentration quenching is likely to account for the low efficiency (2-3%) of the photocurrent generation in photoelectrochemical cells using eosin adsorbed on single crystal TiO₂ electrodes. The advantage of colloidal semiconductor solutions demonstrated in this study is that the undesirable effect of concentration quenching can readily be eliminated and that the efficiency of charge injection is greatly increased by increasing the particle concentration.

An important aspect of this work concerns the fate of the electron photoinjected in the colloidal semiconductor particle, in particular its back reaction with the parent cation. For the first time it has been possible to monitor directly the kinetics of this process. A first-order reaction involving e⁻...EO⁺ pairs associated with the same TiO₂ particle could be clearly discerned from charge recombination via bulk diffusion, and rate constants for these elementary steps have been derived. The encouraging result is that the intraparticle back-electron transfer is ca. 10⁸ times slower than the forward injection. As a consequence, light-induced charge separation can be sustained in the semiconductor particle for at least several microseconds, which is sufficiently long to trap the injected electron by a noble metal deposited on its surface. These results should be important for the understanding of photosensitized electron transfer and photocatalysis which are widely used in photography and artificial photosynthesis.

Acknowledgment. Support of this work by the Schweizerische Nationalfonds zur Förderung der Wissenschaften and CIBA-GEIGY AG, Basel, is gratefully acknowledged. We are grateful to Dr. R. Humphry-Baker for assistance in the light-scattering experiments for particle size determination and to Dr. T. Geiger for performing cyclic voltammetric measurements to obtain the redox potential of eosin in water.

Registry No. TiO₂, 13463-67-7; Eosin, 17372-87-1.

(52) Preliminary experiments show that addition of H₂ does not change the features of the EO⁺ decay in Figure 12a, indicating that \( k_\text{trasp} \) is not affected by H₂. However, hydrogen was used also in the preparation of Pt/TiO₂, and this may have an effect on the type of junction formed.